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                  CAS patent authority coverage expanded
                 ENCOMPLIT/ENCOMPLIT2 search fields enhanced
 NEWS 8
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         APR 28
                 Limits doubled for structure searching in CAS
                  REGISTRY
 NEWS 10 MAY 08 STN Express, Version 8.4, now available
 NEWS 11 MAY 11 STN on the Web enhanced
 NEWS 12 MAY 11 BEILSTEIN substance information now available on
                  STN Easy
                 DGENE, PCTGEN and USGENE enhanced with increased
 NEWS 13 MAY 14
                  limits for exact sequence match searches and
                  introduction of free HIT display format
 NEWS 14
         MAY 15
                 INPADOCDB and INPAFAMDB enhanced with Chinese legal
                  status data
 NEWS 15 MAY 28
                 CAS databases on STN enhanced with NANO super role in
                  records back to 1992
 NEWS 16 JUN 01
                CAS REGISTRY Source of Registration (SR) searching
                  enhanced on STN
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             AND CURRENT DISCOVER FILE IS DATED 06 APRIL 2009.
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-> index bioscience medicine FILE 'DRUGMONOG' ACCESS NOT AUTHORIZED COST IN U.S. DOLLARS

STNCE PILE TOTAL. ENTRY SESSION

FULL ESTIMATED COST INDEX 'ADISCTI, ADISINSIGHT, ADISNEWS, AGRICOLA, ANABSTR, ANTE, ACUALINE, AGUASCI, BIOENG, BIOSIS, BIOTECHABS, BIOTECHOS, BIOTECHNO, CABA, CAPLUS, CEABA-VTB, CIN, CONFSCI, CROPB, CROPU, DDFB, DDFU, DGENE, DISSABS, DRUGB, DRUGMONOG2, DRUGU, EMBAL, EMBASE, ... ENTERED AT 15:35:05 ON 18 JUN 2009

71 FILES IN THE FILE LIST IN STNINDEX

Enter SET DETAIL ON to see search term postings or to view search error messages that display as 0* with SET DETAIL OFF.

-> s (endonucleas? or dnase? or exonucleas? or deoxyribonucleas?) (s) (infect? or diseas? or conditio?) (s) (fung? or albican?)

FILE ADISINSIGHT

10 FILE AGRICOLA

FILE ACUASCI 18 FILE BIOENG

FILE BIOSIS

160 FILE BIOTECHARS

160 FILE BIOTECHDS 12 FILES SEARCHED...

37 FILE BIOTECHNO

63

FILE CABA

8 PILE CAPLUS

FILE CEABA-VTB 747 FILE DGENE

23 FILES SEARCHED...

FILE DISSABS

FILE DRUGU

FILE EMBASE

FILE ESBIOBASE

FILE GENBANK FILE IFIPAT

40 FILE LIFESCI

42 FILES SEARCHED...

FILE MEDLINE

FILE NTIS

49 FILE PASCAL 47 FILES SEARCHED...

FILE PROMT

FILE SCISEARCH

FILE TOXCENTER

FILE USGENE

281 FILE USPATFULL

60 FILES SEARCHED...

38

FILE USPAT2 FILE WPIDS

47 FILE WPINDEX

FILE NLOB

31 FILES HAVE ONE OR MORE ANSWERS, 71 FILES SEARCHED IN STNINDEX

OUE (ENDONUCLEAS? OR DNASE? OR EXONUCLEAS?) OR DEOXYRIBONUCLEAS?) (S) (INFECT ? OR DISEAS? OR CONDITIO?) (S) (FUNG? OR ALBICAN?)

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       9574 GENBANK
F2
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F3
         281 USPATFULL
F4
         160 BTOTECHARS
F5
         160 BIOTECHDS
F6
         63 CABA
F7
          53 ESBIOBASE
F8
          49 LIFESCI
F9
          49 PASCAL
F10
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          47 WPINDEX
          38 HSPAT2
          37 RIOTECHNO
F1.4
          22 TETDAT
F15
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          10 AGRICOLA
F18
          8 CAPLUS
F19
          6 DISSABS
F20
          4 BIOSTS
F21
          4 EMBASE
F22
          4 MEDLINE
          4 MEDLINE
4 SCISEARCH
3 DRUGU
2 CEABA-VTB
1 ADISINSIGHT
1 AQUASCI
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F28
           1 NTIS
F29
           1 PROMT
F30
          1 TOXCENTER
          1 NLDB
F31
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-> file f3-f15 COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 5.44 5.66

FULL ESTIMATED COST

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822 (ENDONUCLEAS? OR DNASE? OR EXONUCLEAS? OR DEOXYRIBONUCLEAS?) (S) (

-> s (endonucleas? or dnase? or exonucleas? or deoxyribonucleas?) (50a) (fung? or albican?)

1401 (ENDONUCLEAS? OR DNASE? OR EXONUCLEAS? OR DEOXYRIBONUCLEAS?) (5CA) (FUNG? OR ALBICAN?)

=> s 13(50a)(infect? or diseas? or conditio?) 6 FILES SEARCHED ... T.4 298 L3 (50A) (INFECT? OR DISEAS? OR CONDITIO?)

-> s 14(s) treatm? L5 64 L4(S) TREATM?

=> dup rem 15 DUPLICATE IS NOT AVAILABLE IN 'USGENE'. ANSWERS FROM THESE FILES WILL BE CONSIDERED UNIQUE PROCESSING COMPLETED FOR L5 L6 58 DUP REM L5 (6 DUPLICATES REMOVED)

=> d ti 15 1-58

ANSWER 1 OF 64 HSPATFHILL on STN HIMAN DNASE IT

ANSWER 2 OF 64 USPATFULL on STN Sensitive detection of bacteria by improved nested polymerase chain reaction targeting the 16S ribosomal RNA gene and identification of bacterial species by amplicon sequencing

ANSWER 3 OF 64 HSPATFHILL on STN TI ANTI-INFECTIVE THERAPY

L5 ANSWER 4 OF 64 USPATFULL on STN

Targeting enzymes of the trna splicing pathway for identification of anti-fungal and/or anti-proliferative molecules

ANSWER 5 OF 64 USPATFULL on STN

TI Methods of identifying compounds that target trna splicing endonuclease and uses of said compounds as anti-fungal agents

ANSWER 6 OF 64 USPATFULL on STN TT Sensitive detection of bacteria by improved nested polymerase chain reaction targeting the 16S ribosomal RNA gene and identification of bacterial species by amplicon sequencing ANSWER 7 OF 64 USPATFULL on STN Compaction assay for assessment of respiratory disease therapy ANSWER 8 OF 64 USPATFULL on STN Nucleic acid probes and methods for detecting clinically important fungal pathogens ANSWER 9 OF 64 USPATFULL on STN Human DNago II ANSWER 10 OF 64 USPATFULL on STN Human DNase ANSWER 11 OF 64 USPATFULL on STN Anti-infective therapy ANSWER 12 OF 64 USPATFULL on STN Immunogenic complex ANSWER 13 OF 64 HISPATFILL OR STN Human DNase II ANSWER 14 OF 64 USPATFULL on STN TΙ 207 human secreted proteins ANSWER 15 OF 64 USPATFULL on STN Compositions and methods for the therapy and diagnosis of colon cancer ANSWER 16 OF 64 USPATFULL on STN Compaction assay for assessment of respiratory disease therapy ANSWER 17 OF 64 USPATFULL on STN Human DNase ANSWER 18 OF 64 USPATFULL on STN Human DNase II L5 ANSWER 19 OF 64 USPATFULL on STN Purified forms of DNase ANSWER 20 OF 64 USPATFULL on STN Compositions and methods for the therapy and diagnosis of pancreatic cancer ANSWER 21 OF 64 USPATFULL on STN DNase Liquid solutions ANSWER 22 OF 64 USPATFULL on STN TI Anti-infective therapy

Compositions and methods for the therapy and diagnosis of colon cancer

ANSWER 23 OF 64 USPATFULL on STN

ANSWER 24 OF 64 USPATFULL on STN

ANSWER 25 OF 64 USPATFULL on STN

Human DNase

TT

- I Compositions and methods for the therapy and diagnosis of ovarian cancer
- 5 ANSWER 26 OF 64 USPATFULL on STN Human DNase
- I Human Dwase
- L5 ANSWER 27 OF 64 USPATFULL on STN TI Purified forms of DNase
- L5 ANSWER 28 OF 64 USPATFULL on STN
- TI Minimizing thermally induced aggregation of DNase in solution with calcium
 - ANSWER 29 OF 64 USPATFULL on STN
- TI Compaction assay for assessment of respiratory disease therapy
- L5 ANSWER 30 OF 64 USPATFULL on STN
- TI Human DNase II
- L5 ANSWER 31 OF 64 HSPATFHILL on STN
- II Gene encoding human Dnase
- 5 ANSWER 32 OF 64 USPATFULL on STN Gene encoding human Dnase
- L5 ANSWER 33 OF 64 HSPATFILL on STN
- TI Purified forms of DNase
- L5 ANSWER 34 OF 64 USPATFULL on STN TI Purified forms of DNASE

in DNA fingerprinting:

- L5 ANSWER 35 OF 64 BIOTECHDS COPYRIGHT 2009 THOMSON REUTERS on STN TI Treating oncological, infectious or somatic diseases comprises acting on extracellular DNA, e.g. circulating in blood plasma using e.g.
 - deoxyribonuclease; liposome-mediated DNA-ase gene transfer and expression in tumor mouse animal model for use in gene therapy
- L5 ANSWER 36 OF 64 BIOTECHDS COPYRIGHT 2009 THOMSON REUTERS on STN
 - Detecting fungi such as Candida albicans or Cryptococcus neoformans, comprises performing nucleic acid amplification using a primer containing an oligonucleotide specific for a base sequences of fungi;
 - Candida albicans, Candida krusei or Cryptococcus neoformans detection using polymerase chain reaction for use in deep-seated mycosis diagnosis and therapy
- 5. ANSWER 37 OF 64 BIOTECHDS COPYRIGHT 2009 THOMSON REUTERS on STN II New Zinc Pinger Protein (2FP) comprising three essential domains useful for diagnosing diseases associated with abnormal genomic structure;
 - transgenic animal and transgenic plant with virus disease-resistance and virus infection therapy and gene therapy in humans
- L5 ANSWER 38 07 64 BIOTECHIS COPYRIGHT 2009 THOMSON REUTERS on STN Simultaneous sequence-specific identification and separation of polyuncleotide fragments, comprises using restriction endonucleases that recomize degenerate bases in their recommitton/cleavage sequence, uneful
 - restriction enzyme, vector expression in host cell, gel electrophoresis and polymerase chain reaction useful disease diagnosis and mutation detection
 - ANSWER 39 OF 64 CABA COPYRIGHT 2009 CABI on STN
 - I Biofilm matrix of Candida albicans and Candida tropicalis: chemical

- composition and role in drug resistance.
- L5 ANSWER 40 OF 64 CABA COPYRIGHT 2009 CABI on STN
 - [Microflora on bean seeds (Phaseolus vulgaris L.)].
 Microflora en semillas de frijol (Phaseolus vulgaris L.).
- LS ANSWER 41 OF 64 CABA COPYRIGHT 2009 CABI on STN
- TI Characterization of a 20 kDa DNase elicitor from Fusarium solani f. sp. phaseoli and its expression at the onset of induced resistance in Pisum sativum.
- L5 ANSWER 42 OF 64 CABA COPYRIGHT 2009 CABI on STN
- TI Transmission of fluconazole-resistant Candida albicans between patients with AIDS and oropharyngeal candidiasis documented by pulsed-field gel electrophoresis.
- L5 ANSWER 43 OF 64 CABA COPYRIGHT 2009 CABI on STN
- TI Investigation of the sequence of colonization and candidemia in nonneutropenic patients.
- L5 ANSWER 44 OF 64 CABA COPYRIGHT 2009 CABI on STN
- TI Studies on watercress chlorotic leaf spot virus and on the control of the fungus vector (Spongospora subterranea) with zinc.
- L5 ANSWER 45 OF 64 Elsevier Biobase COPYRIGHT 2009 Elsevier Science B.V. on
- FI Biofilm matrix of Candida albicans and Candida tropicalis: Chemical composition and role in drug resistance
- L5 ANSWER 46 OF 64 Elsevier Biobase COPYRIGHT 2009 Elsevier Science B.V. on
- TI Identification of medically significant fungal genera by polymerase chain reaction followed by restriction enzyme analysis
- L5 ANSWER 47 OF 64 LIFESCI COPYRIGHT 2009 CSA on STN
- Biofilm matrix of Candida albicans and Candida tropicalis: chemical composition and role in drug resistance
- L5 ANSWER 48 OF 64 LIFESCI COPYRIGHT 2009 CSA on STN
- TI Identification of medically significant fungal genera by polymerase chain reaction followed by restriction enzyme analysis
- L5 ANSWER 49 OF 64 LIFESCI COPYRIGHT 2009 CSA on STN
- TI Transmission of fluconazole-resistant Candida albicans between patients with AIDS and orogharyngeal candidiasis documented by pulsed-field gel electrophoresis
- L5 ANSWER 50 OF 64 PASCAL COPYRIGHT 2009 INIST-CNRS. ALL RIGHTS RESERVED. on STN
- TIEN Biofilm matrix of Candida albicans and Candida tropicalis : chemical composition and role in drug resistance
- L5 ANSWER 51 OF 64 PASCAL COPYRIGHT 2009 INIST-CNRS. ALL RIGHTS RESERVED.
- on STN
 TIEN Transmission of fluconazole-resistant Candida albicans between patients with AIDS and oropharyngeal candidiasis documented by pulsed-field gel electrophoresis
- L5 ANSWER 52 OF 64 WPIDS COPYRIGHT 2009 THOMSON REUTERS on STN TI Identifying compound that modulates fungal tRNA splicing endonuclease activity, involves expressing nucleic acid comprising reporter gene,
 - contacting cell with library of compounds, and detecting expression of

reporter gene

- ANSWER 53 OF 64 WPIDS COPYRIGHT 2009 THOMSON REUTERS on STN System for administration of bloactive compounds by inhalation comprises
- active compound and lipid mixture, useful for delivery of e.g. carboplatin or genes
- ANSWER 54 OF 64 WPIDS COPYRIGHT 2009 THOMSON REUTERS on STN TT Purified forms of non-deamidated and deamidated human DNase - for treatment of pulmonary distress, cystic fibrosis, chronic bronchitis, emphysema, pneumonia, asthma, tuberculosis and fungal infections
- ANSWER 55 OF 64 WPIDS COPYRIGHT 2009 THOMSON RELITERS On STN
- New anti-sense phosphoramidate-linked oligo-nucleotide(s) are more resistant to endo- and exo-nuclease(s) than unmodified phospho-di:ester oliqo-nucleotide(s)
- ANSWER 56 OF 64 USPAT2 on STN
 - Sensitive detection of bacteria by improved nested polymerase chain reaction targeting the 16S ribosomal RNA gene and identification of bacterial species by amplicon sequencing
- ANSWER 57 OF 64 USPAT2 on STN
- Compaction assay for assessment of respiratory disease therapy
- ANSWER 58 OF 64 USPAT2 on STN
- Human DNase
- -> d ibib abs 15 1 3 11 17 19 21 31 35 39 42 54 57
- L5 ANSWER 1 OF 64 USPATFULL on STN ACCESSION NUMBER: 2009:45775 USPATFULL
- TITLE: HUMAN DNASE II
- INVENTOR(S): Baker, Kevin P., Darnestown, MD, UNITED STATES Baron, Will F., Moorpark, CA, UNITED STATES
- PATENT ASSIGNEE(S): Genentech, Inc., South San Francisco, CA, UNITED STATES (U.S. corporation) NUMBER KIND DATE

PATENT INFORMATION:	US 20090041742 A1 20090212
APPLICATION INFO.:	US 2007-740860 A1 20070426 (11)
RELATED APPLN. INFO.:	Continuation of Ser. No. US 2004-990207, filed on 15
	Nov 2004, ABANDONED Continuation of Ser. No. US
	2003-408167, filed on 4 Apr 2003, ABANDONED
	Continuation of Ser. No. US 2001-861034, filed on 18
	May 2001, Pat. No. US 6569429 Division of Ser. No. US
	1996-639294, filed on 25 Apr 1996, Pat. No. US 6265195
DOCUMENT TYPE:	Utility
FILE SEGMENT:	APPLICATION
LEGAL REPRESENTATIVE:	GENENTECH, INC., 1 DNA WAY, SOUTH SAN FRANCISCO, CA,

94080, US

NUMBER OF CLAIMS: 16 EXEMPLARY CLAIM: NUMBER OF DRAWINGS: 2 Drawing Page(s)

LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention relates to a novel human deoxyribonuclease, referred to as human DNase II. The invention provides nucleic acid sequences

encoding human DNase II, thereby enabling the production of human DNase II by recombinant DNA methods in quantities sufficient for clinical use. The invention also relates to pharmaceutical compositions and diagnostic and therapeutic uses of human DNase II.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 3 OF 64 USPATFULL on STN ACCESSION NUMBER: 2008:30136 USPATFULL

TITLE: ANTI-INFECTIVE THERAPY

INVENTOR(S): Shak, Steven, Burlingame, CA, UNITED STATES

PATENT ASSIGNEE(S): Genentech, Inc., South San Francisco, CA, UNITED STATES (U.S. corporation)

(U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 20080026426 A1 20080131

PATENT INFORMATION: US 20080026426 Al 20080131 APPLICATION INFO: US 2007-862934 Al 20070927 (11) RELATED APPLN. INFO: Continuation of Ser. No. US 2004-839046, filed on 4 May

2004. GRANTED, Pat. No. US 2009-053900, filed on a may 2004. GRANTED, Pat. No. US 7297526 Continuation of Ser. No. US 2001-5675, filed on 7 Nov 2001, ABANDONED Continuation of Ser. No. US 2000-669306, filed on 25

Sep 2000, ABANDONED Continuation of Ser. No. US 1996-761578, filed on 9 Dec 1996, ABANDONED

Continuation of Ser. No. US 1995-528876, filed on 15 Sep 1995, ABANDONED Continuation of Ser. No. US 1993-117584, filed on 3 Sep 1993, ABANDONED Division of

1993-117584, filed on 3 Sep 1993, ABANDONED Division Ser. No. US 1992-914226, filed on 13 Jul 1992, ABANDONED Continuation of Ser. No. US 1989-448038,

filed on 8 Dec 1989, ABANDONED Continuation-in-part of Ser. No. US 1988-289958, filed on 23 Dec 1988,

ABANDONED DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION
LEGAL REPRESENTATIVE: GENERATECH, INC., 1 DNA WAY, SOUTH SAN FRANCISCO, CA,

LEGAL REPRESENTATIVE: GENERATECH, 1 94080, US

PATENT ASSIGNEE(S):

NUMBER OF CLAIMS: 13 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 18 Drawing Page(s) LINE COUNT: 1866

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB DNA isolates coding for human DNase and methods of obtaining such DNA are provided, together with expression systems for recombinant

production of human DNase useful in therapeutic or diagnostic compositions.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 11 OF 64 USPATFULL on STN ACCESSION NUMBER: 2005:10922 USPATFULL

ACCESSION NUMBER: 2005:10922 USPATFULL TITLE: Anti-infective therapy

TITLE: Anti-infective therapy
INVENTOR(S): Shak, Steven, Burlingame, CA, UNITED STATES

Genentech, Inc. (U.S. corporation)

PATENT INFORMATION: US 20050009056 A1 20050113 US 7297526 B2 20071120 APPLICATION INFO:: US 2004-039046 A1 20040504 (10)

RELATED APPLN. INFO.: Continuation of Ser. No. US 2001-5675, filed on 7 Nov

2001, ABANDONED Continuation of Ser. No. US 2000-669306, filed on 25 Sep 2000, ABANDONED

Continuation of Ser. No. US 1996-761578, filed on 9 Dec 1996, ABANDONED Continuation of Ser. No. US 1995-528876, filed on 15 Sep 1995, ABANDONED Continuation of Ser. No. US 1993-117584, filed on 3 Sep 1993, ABANDONED Division of Ser. No. US 1992-94726, filed on 13 Jul 1992, ABANDONED Continuation of Ser. No. 15 1998-428958, filed on 6 Dec 1999, ABANDONED Continuation of Ser. No. 15 1989-249586, filed on 25 Dec 1998.

DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: GENERATECH, INC., 1 DNA WAY, SOUTH SAN FRANCISCO, CA,

NUMBER OF CLAIMS: 35 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 18 Drawing Page(s)

LINE COUNT: 1917
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

DNA isolates coding for human DNase and methods of obtaining such DNA are provided, together with expression systems for recombinant production of human DNase useful in therapeutic or diagnostic

compositions.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 17 OF 64 USPATFULL on STN

ACCESSION NUMBER: 2003:142950 USPATFULL TITLE: Human DNase

INVENTOR(S): Rosen, Craig A., Laytonsville, MD, United States
Ruben, Steven M., Olney, MD, United States

PATENT ASSIGNEE(S): Adams, Mark D., North Potomac, MD, United States
Human Genome Sciences, Inc., Rockville, MD, United
States (U.S. corporation)

RELATED APPLN. INFO.: Division of Ser. No. US 1998-54989, filed on 3 Apr 1998, now patented, Fat. No. US 6251468, issued on 16 Jun 2001 Division of Ser. No. US 1995-468012, filed on 6 Jun 1995, now reatented, Fat. No. US 5830744, issued

on 3 Nov 1998 Continuation-in-part of Ser. No. WO 1994-US4954, filed on 5 May 1994

DOCUMENT TYPE: Utility
FILE SEGMENT: GRANTED
PRIMARY EXAMINER: Saidha, Tekchand
ASSISTANT EXAMINER: Walicka, Malgorrata A.

LEGAL REPRESENTATIVE: Human Genome Sciences, Inc. NUMBER OF CLAIMS: 48
EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 6 Drawing Figure(s); 6 Drawing Page(s)

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

One nucleation. Defice replaying the set DRS (RMA) encoding such polyopetide and a procedure for protecting such polyopetide by recombinant techniques is disclosed. Also disclosed are methods for utilizing such polyopetide of proventing and/or treating bronchopsinomary conditions. Diagnostic analys for identifying mutations in nucleic acid sequence encoding a party of the property of the propert

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 19 OF 64 USPATFULL on STN

ACCESSION NUMBER: 2003:112537 USPATFULL Purified forms of DNase

INVENTOR(S): Frenz, John, Millbrae, CA, UNITED STATES

Shire, Steven J., Belmont, CA, UNITED STATES

Sliwkowski, Mary B., San Carlos, CA, UNITED STATES PATENT ASSIGNEE(S): Genentech, Inc. (U.S. corporation)

NUMBER KIND DATE PATENT INFORMATION: IIS 20030077267 Α1 20030424 IIS 6932965 R2 20050823

APPLICATION INFO . . IIS 2002-155407 21 20020522 (10)

RELATED APPLN. INFO.: Continuation of Ser. No. US 2000-638112, filed on 11 Aug 2000, GRANTED, Pat. No. US 6440412 Continuation of

Ser. No. US 1997-942561, filed on 1 Oct 1997, ABANDONED Continuation of Ser. No. US 1996-634125, filed on 19 Apr 1996, ABANDONED Continuation of Ser. No. US

1995-409631, filed on 22 Mar 1995, ABANDONED Continuation of Ser. No. US 1994-348284, filed on 30

Nov 1994, ABANDONED Continuation of Ser. No. US

1993-116186, filed on 2 Sep 1993, ABANDONED

Continuation of Ser. No. US 1992-895300, filed on 8 Jun 1992, GRANTED, Pat. No. US 5279823

DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION GENENTECH, INC., 1 DNA WAY, SOUTH SAN FRANCISCO, CA, LEGAL REPRESENTATIVE:

94080 NUMBER OF CLAIMS: 19

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

10 Drawing Page(s) LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT. The present invention provides the identification and characterization aB

of two components of a recombinant preparation of DNase. These components are the purified deamidated and non-deamidated human DNases. Taught herein are the separation of these components and the use of the non-deamidated species as a pharmaceutical per se, and in particular in compositions wherein the species is disclosed within a plastic vial, for use in administering to patients suffering from pulmonary distress.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 21 OF 64 USPATFULL on STN ACCESSION NUMBER: 2003:78612 USPATFULL

DNase Liquid solutions INVENTOR(S):

Chan, Hak-Kim, North Sydney, AUSTRALIA Gonda, Igor, San Francisco, CA, UNITED STATES Shire, Steven J., Belmont, CA, UNITED STATES

Weck, Suzanne Sin-Mui Lo, Mountain View, CA, UNITED STATES

PATENT ASSIGNEE(S): Genentech, Inc. (non-U.S. corporation)

NUMBER KIND DATE US 20030054532 A1 20030320 PATENT INFORMATION: IIS 7018825 B2 20060328

APPLICATION INFO .: US 2002-76213 A1 20020212 (10) RELATED APPLN. INFO.: Continuation of Ser. No. US 1996-696955, filed on 3 Dec

1996, GRANTED, Pat. No. US 6383788 A 371 of

International Ser. No. WO 1995-US2457, filed on 28 Feb 1995, PENDING A 371 of International Ser. No. US

1995-377527, filed on 20 Jan 1995, ABANDONED Continuation of Ser. No. US 1994-206504, filed on 4 Mar

1994, ABANDONED

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: Richard F. Trecartin, Esq., FLEHR HOHBACH TEST

ALBRITTON & HERBERT LLP, Four Embarcadero Center, Suite 3400, San Francisco, CA, 94111-4187

NUMBER OF CLAIMS: 16 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 8 Drawing Page(s)

LINE COUNT: 991
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention relates to the use of calcium ion and/or sugars to ministre thermal aggregation of DNase and to the use of calcium ion to stabilize liquid solutions of DNase, the solutions having a pH of less than neutral. DNase is the active pharmaceutical principle and the solutions may contain other pharmaceutically acceptable exciplents making them suitable for pharmaceutical administration. In the fination intanno, solicium ion/supar ministrate the effects of thermal aggregation of the pharmaceutical process of the second pharmaceutical pharmaceutical pharmaceutical all solutions from protein precipitation.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L5 ANSWER 31 OF 64 USPATFULL on STN

ACCESSION NUMBER: 2001:97673 USPATFULL TITLE: Gene encoding human Dnase

INVENTOR(S): Rosen, Craig, Laytonsville, MD, United States

Ruben, Steven M., Olney, MD, United States Adams, Mark D., North Potomac, MD, United States

PATENT ASSIGNEE(S): Human Genome Sciences, Inc., Rockville, MD, United States (U.S. corporation)

APPLICATION INFO.: US 1998-54989 19980403 (9)
RELATED APPLN. INFO.: Division of Ser. No. US 468012, now patented, Pat. No. US 5830744

DOCUMENT TYPE: Utility
FILE SEGMENT: GRANTED
PRIMARY EXAMINER: Stole, Einar

LEGAL REPRESENTATIVE: Human Genome Sciences, Inc.

NUMBER OF CLAIMS: 51 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 7 Drawing Figure(s); 7 Drawing Page(s) LINE COUNT: 1273

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A human DMase polypeptide and DMA (BMA) encoding such polypeptide and a procedure for producing such polypeptide by recombinant techniques is disclosed. Also disclosed are methods for utilizing such polypeptide for preventing and/or treating bronchopulnonary conditions. Diagnostic assays for identifying mutations in nucleic acid sequence encoding a

polypeptide of the present invention and for detecting altered levels of the polypeptide of the present invention are also disclosed.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

TITLE:

L5 ANSWER 35 OF 64 BIOTECHDS COPYRIGHT 2009 THOMSON REUTERS on STN ACCESSION NUMBER: 2005-07193 BIOTECHDS

Treating oncological, infectious or somatic diseases

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comprises acting on extracellular DNA, e.g. circulating in
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blood plasma using e.g. deoxyribonuclease;

liposome-mediated DNA-ase gene transfer and expression in

tumor mouse animal model for use in gene therapy AUTHOR: TETS V V; GENKIN D D; TETS G V

PATENT ASSIGNEE: TETS V V; GENKIN D D
PATENT INPO: WO 2005007187 27 Jan 2005

APPLICATION INFO: WO 2003-RU304 14 Jul 2003

PRIORITY INFO: WO 2003-304 14 Jul 2003; WO 2003-304 14 Jul 2003

DOCUMENT TYPE: Patent

LANGUAGE: Unavailable RS OTHER SOURCE: WFI: 2005-132270 [14]

AN 2005-07193 BIOTECHDS

AB DERWENT ABSTRACT:

NOVELTY - Treating oncological, infectious or somatic diseases comprises acting on extracellular DNA, e.g. circulating in blood plasma.

acting on water bases and bases of the second of the secon

efficacy of the treatment of oncological, infectious or somatic diseases by monitoring the amount, molecular weight, fraction ratio,

protein, lipid and sugar binding and/or nucleotide sequence of DNA freely circulating in blood plasma; (3) use of blood plasma DNA and

extracellular microbial DNA to detect DNA involved in the onset and development of diseases, comprising cloning, sequencing and identifying genes, unique sequences and repeat sequences for subsequent

study. ACTIVITY - Cytostatic; Antibacterial; Fungicide; Protozoacide. Mice with transplanted Ehrlich tumors were treated twice a

day on days 3-7 post transplantation by intraperitoneal injection with DNase I (1 mg/kg) in phosphate buffer (200 mul). Tumor volume on day 7 was reduced by 61 % compared with controls.

MECHANISM OF ACTION - Extracellular DNA inactivator. USE - Treating oncological, infectious or somatic diseases, including malignant tumors, bacterial, fungal

or protozoal infections, noninfectious somatic diseases and diseases caused by the accumulation of somatic mutations. (96 pages)

L5 ANSWER 39 OF 64 CABA COPYRIGHT 2009 CABI on STN

ACCESSION NUMBER: 2006:205112 CABA DOCUMENT NUMBER: 20063192996

DOCUMENT NUMBER: 20063192996
TITLE: Biofilm matrix of Candida albicans and Candida

tropicalis: chemical composition and role in drug

AUTHOR: Al-Fattani, M. A.; Douglas, L. J.

CORPORATE SOURCE: Division of Infection and Immunity, Institute of Biomedical and Life Sciences, Joseph Black Building, University of Glasqow, Glasqow Glasqow and Seq. UK.

J.Douglas@bio.gla.ac.uk

SOURCE: Journal of Medical Microbiology, (2006) Vol. 55, No. 8, pp. 999-1008, 44 ref.

Publisher: Society for General Microbiology. Reading ISSN: 0022-2615

URL: www.sgm.ac.uk

DOI: 10.1099/jmm.0.46569-0 PUB. COUNTRY: United Kingdom

DOCUMENT TYPE: Journal LANGUAGE: English

LANGUAGE: English
ENTRY DATE: Entered STN: 6 Dec 2006

Last Updated on STN: 6 Dec 2006

AR. Matrix material was extracted from biofilms of Candida albicans and Candida tropicalis and analysed chemically, Both preparations contained carbohydrate, protein, hexosamine, phosphorus and uronic acid. However, the major component in C. albicans matrix was glucose (32%), whereas in C. tropicalis matrix it was hexosamine (27%). Biofilms of C. albicans were more easily detached from plastic surfaces by treatment with the enzyme lyticase ([beta]-1.3-glucanase) than were those of C. tropicalis, Biofilms of C. albicans were also partially detached by treatment with proteinase K, chitinase, DNase I, or [beta]-N-acetylqlucosaminidase, whereas C. tropicalis biofilms were only affected by lipase type VII or chitinase. To investigate a possible role for the matrix in biofilm resistance to antifungal agents, biofilms of C. albicans were grown under conditions of continuous flow in a modified Robbins device (MRD). These biofilms produced more matrix material than those grown statically, and were significantly more resistant to amphotericin B. Biofilms of C. tropicalis synthesized large amounts of matrix material even when grown statically, and such biofilms were completely resistant to both amphotericin B and fluconazole. Mixed-species biofilms of C. albicans and a slime-producing strain of Staphylococcus epidermidis (RP62A), when grown statically or in the MRD, were also completely resistant to amphotericin B and fluconazole. Mixed-species biofilms of C. albicans and a slime-negative mutant of S. epidermidis (M7), on the other hand, were completely drug resistant only when grown under flow conditions. These results demonstrate that the matrix can make a significant contribution to drug resistance in Candida biofilms, especially under conditions similar to those found in catheter infections in vivo, and that the composition of the matrix material is an important determinant in resistance.

L5 ANSWER 42 OF 64 CABA COPYRIGHT 2009 CABI on STN ACCESSION NUMBER: 96:54354 CABA

DOCUMENT NUMBER:

19962000515 TITLE:

Transmission of fluconazole-resistant Candida albicans between patients with AIDS and

oropharyngeal candidiasis documented by pulsed-field gel electrophoresis

Barchiesi, F.; Hollis, R. J.; Poeta, M. del;

McGough, D. A.; Scalise, G.; Rinaldi, M. G.; Pfaller, M. A.; Del Poeta, M.

CORPORATE SOURCE: Pungus Testing Laboratory, Department of Pathology, University of Texas Health Science Center at San Antonio, 7703 Floyd Curl Drive, San Antonio, Texas 78284-7750. USA.

SOURCE: Clinical Infectious Diseases, (1995) Vol. 21, No. 3, pp. 561-564. 25 ref.

ISSN: 1058-4838

DOCUMENT TYPE: Journal LANGUAGE: English

ENTRY DATE: Entered STN: 30 Apr 1996 Last Updated on STN: 30 Apr 1996

Electrophoretic karyotype and restriction endonuclease analysis of genomic DNA were used to type 9 isolates of C. albicans from the oral cavities of 2 AIDS patients (husband and wife, aged 38 and 34 vr) from Texas, USA, who had infections that had become resistant to fluconazole treatment (400 mg/d). The in vitro susceptibilities of sequential isolates to fluconazole, itraconazole and investigational drug D0870 were also evaluated. DNA analysis showed that the isolates responsible for fluconazole-resistant episodes of oropharyngeal candidosis in the 2 patients were genetically related. In vitro susceptibility to fluconazole correlated well with clinical outcome. Although the min. inhibitory concn of itraconazole and D0870 for fluconazole-resistant isolates were higher than those for fluconazole-susceptible isolates. Both itraconazole and D0870 showed good in vitro activity against the isolates tested.

L5 ANSWER 54 OF 64 WPIDS COPYRIGHT 2009 ACCESSION NUMBER: 1994-007528 [01] WPIDS DOC. NO. CPI: C1994-003051 [01]

THOMSON REUTERS on STN

Purified forms of non-deamidated and deamidated human DNase - for treatment of pulmonary distress, cystic fibrosis, chronic bronchitis, emphysema, pneumonia, asthma, tuberculosis and fungal infections

DERWENT CLASS: B04; C06; D16 INVENTOR: FRENZ J; FRENZ

FRENZ J; FRENZ J H; FRENZ J M; SHIRE S; SHIRE S J; SILIWKOWSKI M B; SLIWKOWSKI M B; SLIWOWSKI M B

PATENT ASSIGNEE: (GETH-C) GENENTECH INC COUNTRY COUNT: 43

PATENT INFO ABBR.:

TITLE:

PAT	ENT NO	KIN	DATE	MEEK	LA	PG	MAIN I	PC
	9325670	A1		(199401)*		38[9]		
	5279823	A	19940118	(199404)	EN	24[9]		
ΑU			19940104	(199417)	EN			
	9405549	A	19941125	(199508)	FI			
NO	9404752	A	19941208	(199510)	NO			
ZA			19950125	(199510)	EN	39		
			19950329		EN			
	644932	A1	19950329	(199517)	EN			
	9403032	A3	19950614	(199532)	CS			
DE	4392749	T	19950824	(199539)	DE	[0]		
	07507455	56	19950824	(199539) (199542) (199615)	JA	16[0]		
SK	9401495	A3	19960110	(199615)	SK			
	2282140		19960417	(199619)	EN			
NZ	253559	A	19961126	(199701)	EN			
IL	105724	A	19970610	(199730)	EN			
ΗU	70468	T	19951030	(199732)	HU			
ΑU	682822	В	19971023	(199750)	EN			
US	70468 682822 5783433 9306670 1013284	A	19980721	(199836)	EN			
BR	9306670	A	19981208	(199903)	PT			
EP	1013284	A2	20000628	(200035)	EN			
ΕP	644932	B1	20000809	(200039)	EN			
NZ	299257	A	20000825	(200049)	EN			
DE	69329200	Е	20000914	(200053)	DE			
ES	2150447	T3	20001201	(200105)	ES			
	219549	В	20010528	(200140)	HU			
RO	117188	B1	20011130	(200225)	RO			
KR	302092	В	20011022	(200236)	KO			
	323357	В	20010528 20011130 20011022 20020219	(200257)	KO			
	6440412	B1	20020827	(200259)	EN			
SK	282957	B6	20030109	(200309)	SK			
JP	3383307	B2	20030304	(200319)	JA	28		
US	20030077267	A1	20030424	(200330)	EN			
CZ	293105	B6	20040218	(200430)	CS			
CA	2137237 2238320 318644	C	20041026	(200471)	EN			
RU	2238320	C2	20041020	(200476)	RU			
NO	318644	B1	20050425	(200530)	NO			
	6932965	B2	20050823	(200556)	EN			

APPLICATION DETAILS:

PATENT NO KIND	APPLICATION	DATE
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EP 1	013284 A2		2000-101817 19930528
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	323357 B		2001-703922 20010328
	20030077267 A1		2002-155407 20020522
US 6	5932965 B2	US	2002-155407 20020522

FILING DETAILS:

PA:	TENT NO	KIND		PA1	TENT NO	
AU	682822 B		Previous Publ	AU		1 A
CZ	293105 B6		Previous Publ	CZ	9403032	2 A
EP	644932 B1		Related to	EP	101328	4 A
EP	1013284 A2		Div ex	EP	644932	A
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ES	2150447 T3		Based on	EP	644932	A
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	282957 B6		Previous Publ	SK	940149	5 A
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WO 9325670 A
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     CZ 293105 B6
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     RU 2238320 C2
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PRIORITY APPLN. INFO: US 1992-895300
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                                          19971001
                     US 2000-638112
                                         20000811
                     US 2002-155407
                                         20020522
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AN 1994-007528 [01] WPIDS AB WO 1993025670 A1 UPAB: 20060108

Process (I) comprises separating deamidated and non-deamidated human DNase I from a mixture.

Also claimed are: (1) purified demaindated human DMassy (2) purified non-claimed are: (1) a plaramaceutical compon. consisting of the doubt distributed human DMassaceutically acceptable excipient; deducted human DMassaceutically acceptable excipient; or components of the demain acceptable of the desired product of the demain demain acceptable excipient (5) a pharamaceutically acceptable excipient; (5) a pharamaceutical components of the demain de

greater than 3 weeks. Separation process requires a tentacle cation exchange resin, an immobilised heparin resin and an immobilised non-hydrolysable DNA analogue resin.

USE/ADVANTAGE - beamidated human (Mass or non-deamidated human (Mass or may be used for treatment of patients having an accumulation of viscous, NON-containing material. Admin. of purified (Masses pref. is via directly into the airway passages for the treatment of patients having pulsonary diseases e.g. chronic bronchitis, cystic patients having pulsonary diseases e.g. chronic bronchitis, cystic plotosis or emphysems. The Masses are employed for enzymatic alteration of the viscoelasticity of muoous, for treatment of infectious rememonis, authon, tuberculosis and funnal.

infectious pneumonia, asthma, tuberculosis and fungal mises and infections. — In an example, purfiled deamidated humans mises and infections. — In an example, purfile deamidated human way were prepared by TCX chromatography. NMsse enzymatic activity, synthetic double stranded NRA, 25 hase pairs in length, was labelled with dimitrophenol (NRF) on one end and with biotin on the other end. Bydrolysis of the substrate by DNsse was detected by cogitive of the reaction product on microtiture plate wells streptavidin-horsexadish peroxidases. Specific activity of stability attempts of the samples was correlated (r power (2) = 0.613; m-5) with the extent of IMsse deamidation (range 27% = 53%). Extrapolation of the least squares linear equation provided an extinate that the specific activity of deamidated

Member (0002)

AREO US 5279823 A HPAR 20060108

Deamidated and non-deamidated human DNases have been prepd. by recombinant DNA methods and sepal, and purified by chromatography with a heparin or non-hydrolysable DNA manlogue bonded to a resin or other support medium as adsorbent. These enzymes are phosphordiesterases that cleave polydeoxyribonucleic acids. Pharmacoutical compans, contq. deamidated or

non-deamidated DNase and the usual carriers and additives reduce the viscoelasticity of pulmonary secretions.

USE/ADVANTAGE - The prods. are therapeutics for chronic bronchitis, cystic fibrosis, emphysema, etc. The recombinant enzymes are free from contamination with proteases and other proteins.

Member (0006)

ABEQ ZA 9303724 A UPAB 20060108

Process (1) comprises segs. desaidated and non-desaidated human DNase I from a maker. (1) partified desaidated human DNase (2) partified in the process of t

of human DNase by preparing a compan. comprising non-deam dated human DNase in an aq. soin. having pH 4.5 - 6.8 and storing the compan. for greater than 3 weeks.

Spon. process requires a tentacle cation exchange resin, an immobilised non-baytrnly-sable DNA analogue resin.

heparin remin and an immebilized non-hydrolymable DMA analogue remin. UBS./ADVANTAGE. Demaidated human DMase or non-demaidated human DMase and the property of the property

alteration of the viscoelasticity of macous, for treatment of patients with abnormal viscous, pursuents sceretions e.g. with infectious permaonis, asthms, tweetermines and fungal purified non-deanidated human Obase for use in this atty were proped by TCX chromatography. Obase enzymatic activity, synthetic double stranded MDA, 25 hase pairs in langth, wer labelled with distrepence (ORP) on one

was detected by capture of the reaction prode, on microtiter plate well-tocoated with antibody to BNP and by quantitation of the intact probe with streptavidin-horseradish peroxidase. Specific activity of stability samples was correlated (r power [2] - 0.613; n=5) with the extent of DNsae deamidation (range 27% - 53%). Extrapolation of the least squares linear human DNsae was approx. 77% lower than that of non-deamidated human DNsae.

Member (0010) ABEO DE 4392749 T UPAB 20060108

Process (I) comprises sepg. deamidated and non-deamidated human DNase I from a mixt..

Firm a subt. Arrs: (1) purified desaidated human DMsser (2) purified desaidated human DMsser (3) purified desaidated human DMsser (3) a pharmacoutical compan, consisting of the desaidated human DMsser and opt. a pharmacoutically acceptable excipient; (4) a pharmacoutically acceptable excipient; (5) a pharmacoutical compan comprising non-desaidated human DMsser and a plastic vial, and (6) atomoscopising non-desaidated human DMsser and a plastic vial, and (6) atomoscopising non-desaidated human DMsser and plastic vial, and (6) atomoscopising non-desaidated human DMsser and subt. A plastic vial of the compan. For greater than 3 work, huming pH 4.5 – 6.8 and storing the compan, for greater than 3 work, huming pH 4.5 – 6.8 and storing the compan, for

Sepn. process requires a tentacle cation exchange resin, an immobilized heparin resin and an immobilized non-hydrolysable DNA analogue resin. USS/ADVANYAGE - Deamidated human DNase or non-deamidated human DNase may be used for treatment of patients having an accumulation of

viscous, DNA-contg. material. Admin. of purified DNases pref. is via

direct inhalation into the lungs. Non-deamidated human DNase may be admin. directly into the airway passages for the treatment of patients having pulmonary diseases e.g. chronic bronchitis, cystic fibrosis or emphysema. The DNases are employed for enzymatic alteration of the viscoelasticity of mucous, for treatment of patients with abnormal viscous, purulent secretions e.g. with infectious pneumonia, asthma, tuberculosis and fungal infections. - In an example, purified deamidated human DNase and purified non-deamidated human DNase for use in this study were prepd, by TCX chromatography. DNase enzymatic activity, synthetic double stranded DNA, 25 base pairs in length, was labelled with dimitrophenol (DNP) on one end and with biotin on the other end. Hydrolysis of the substrate by DNase was detected by capture of the reaction prods. on microtiter plate wells coated with antibody to DNP and by quantitation of the intact probe with streptavidin-horseradish peroxidase. Specific activity of stability samples was correlated (r power(2) = 0.613; n=5) with the extent of DNase deamidation (range 27% - 93%). Extrapolation of the least squares linear equation provided an estimate that the specific activity of deamidated human DNase was approx. 77% lower than that of non-deamidated human DNase.

Member (0011) ABEO JP 07507455 W UPAB 20060108

Process (I) comprises sepg. deamidated and non-deamidated human DNase I from a mixt...

Also claimed are: (1) purified deamidated human DNase; (2) purified non-deamidated human DNase; (3) a pharmaceutical compsn. consisting of the deamidated human DNase and opt, a pharmaceutically acceptable excipient; (4) a pharmaceutical compsn. consisting on non-deamidated human DNase and opt. a pharmaceutically acceptable excipient; (5) a pharmaceutical compen. comprising non-deamidated human DNase in a plastic vial; and (6) storage of human DNase by preparing a compsn. comprising non-deamidated human DNase in an ag. soln. having pH 4.5 - 6.8 and storing the compsn. for greater than 3 weeks.

Sepn. process requires a tentacle cation exchange resin, an immobilised beparin resin and an immobilised non-bydrolysable DNA analogue resin. USE/ADVANTAGE - Deamidated human DNase or non-deamidated human DNase

may be used for treatment of patients having an accumulation of

viscous, DNA-contq. material. Admin. of purified DNases pref. is via direct inhalation into the lungs. Non-deamidated human DNase may be admin. directly into the airway passages for the treatment of patients having pulmonary diseases e.g. chronic bronchitis, cystic fibrosis or emphysema. The DNases are employed for enzymatic alteration of the viscoelasticity of mucous, for treatment of patients with abnormal viscous, purulent secretions e.g. with infectious pneumonia, asthma, tuberculosis and fungal infections. - In an example, purified deamidated human DNase and purified non-deamidated human DNase for use in this study were prepd, by TCX chromatography. DNase enzymatic activity, synthetic double stranded DNA, 25 base pairs in length, was labelled with dimitrophenol (DNP) on one end and with biotin on the other end. Hydrolysis of the substrate by DNase was detected by capture of the reaction prods. on microtiter plate wells coated with antibody to DNP and by quantitation of the intact probe with streptavidin-horseradish peroxidase. Specific activity of stability samples was correlated (r power(2) = 0.613; n=5) with the extent of DNase deamidation (range 27% - 93%). Extrapolation of the least squares linear equation provided an estimate that the specific activity of deamidated

Member (0018) ABEO US 5783433 A UPAB 20060108

human DNase was approx. 77% lower than that of non-deamidated human DNase. Process (I) comprises sepg. deamidated and non-deamidated human DNase I from a mixt...

Also claimed arcs: (1) purified desmidated human DNase; (2) purified non-desmidated human DNase; (3) a pharmaceutical compon. consisting of the desmidated human DNase and opt. a pharmaceutically acceptable excipient; (4) a pharmaceutically acceptable excipient; (5) a pharmaceutical compon. compising non-desmidated human DNase and opt. a pharmaceutically acceptable excipient; (5) a pharmaceutical component of the pharmaceutical component and pharmaceutically acceptable acceptaint on a plantic vial, and (6) storage of human DNase in praparing a compon. comprising non-desmidated human grant DNase by greatly as compon. comprising non-desmidated human grant DNase by greatly as a second property of the compon. for greatly than 3 weeks.

Sepn. process requires a tentacle cation exchange resin, an immobilised heparin resin and an immobilised non-hydrolysable DNA analogue resin.

USLADVANTAGE - Deamidated human DNase or non-deamidated human DNase

may be used for treatment of patients having an accumulation of viscous, DNA-contg, material, Admin, of purified DNases pref, is via direct inhalation into the lungs. Non-deamidated human DNase may be admin. directly into the airway passages for the treatment of patients having pulmonary diseases e.g. chronic bronchitis, cystic fibrosis or emphysema. The DNases are employed for enzymatic alteration of the viscoelasticity of mucous, for treatment of patients with abnormal viscous, purulent secretions e.g. with infectious pneumonia, asthma, tuberculosis and fungal infections. - In an example, purified deamidated human DNase and purified non-deamidated human DNase for use in this study were prend, by TCX chromatography. DNase enzymatic activity, synthetic double stranded DNA, 25 base pairs in length, was labelled with dinitrophenol (DNP) on one end and with biotin on the other end. Hydrolysis of the substrate by DNase was detected by capture of the reaction prods. on microtiter plate wells coated with antibody to DNP and by quantitation of the intact probe with streptavidin-horseradish peroxidase. Specific activity of stability samples was correlated (r power(2) - 0.613; n-5) with the extent of DNase deamidation (range 27% - 93%). Extrapolation of the least squares linear equation provided an estimate that the specific activity of deamidated human DNase was approx. 77% lower than that of non-deamidated human DNase.

Member (0020)

ABEQ EP 1013284 A2 UPAB 20060108

Process (I) comprises sepg. deamldated and non-deamldated human DNase I True a slake. There is a large than 1 the second to the

Sepn. process requires a tentacle cation exchange resin, an immobilised heparin resin and an immobilised non-hydrolysable DNA analogue resin. USE/ADVANTAGE — Deamidated human DNase or non-deamidated human DNase

may be used for treatment of patients having an accumulation of viscous, RNA-contp, material. Admin. of purified DNases pref. is via direct inhalation into the lungs. Non-deamidated human DNase may be admin. directly into the airway pasages for the treatment of patients having pulmonary diseases e.g. chronic bronchitis, cystic fibrosis or emphysens. The DNases are employed for enzymatic

alteration of the viscoelasticity of mucous, for treatment of patients with abnormal viscous, purulent secretions e.g. with infectious pneumonia, asthma, tuberculosis and fungal

infections. - In an example, purified deamidated human DNase and purified non-deamidated human DNase for use in this study were prepd. by TCX chromatography. DNase enzymatic activity, synthetic double stranded

DNA, 25 base pairs in length, was labelled with dinitrophenol (DNP) on one end and with biotin on the other end. Hydrolysis of the substrate by DNase was detected by capture of the reaction prods, on microtiter plate wells coated with antibody to DNP and by quantitation of the intact probe with streptavidin-horseradish peroxidase. Specific activity of stability samples was correlated (r power(2) - 0.613; n-5) with the extent of DNase deamidation (range 27% - 93%). Extrapolation of the least squares linear equation provided an estimate that the specific activity of deamidated human DNase was approx. 77% lower than that of non-deamidated human DNase.

Member (0021) ABEQ EP 644932 B1 HPAR 20060108

Process (I) comprises sepg. deamidated and non-deamidated human DNase I

from a mixt

Also claimed are: (1) purified deamidated human DNase: (2) purified non-deamidated human DNase; (3) a pharmaceutical compsn. consisting of the deamidated human DNase and opt. a pharmaceutically acceptable excipient; (4) a pharmaceutical compsn. consisting on non-deamidated human DNase and opt. a pharmaceutically acceptable excipient; (5) a pharmaceutical compsn. comprising non-deamidated human DNase in a plastic vial; and (6) storage of human DNase by preparing a compsn. comprising non-deamidated human DNase in an ag, soln, having pH 4.5 - 6.8 and storing the compan, for greater than 3 weeks.

Sepn. process requires a tentacle cation exchange resin, an immobilised heparin resin and an immobilised non-hydrolysable DNA analogue resin. USE/ADVANTAGE - Deamidated human DNase or non-deamidated human DNase

may be used for treatment of patients having an accumulation of viscous, DNA-contg. material. Admin. of purified DNases pref. is via direct inhalation into the lungs. Non-deamidated human DNase may be admin. directly into the airway passages for the treatment of patients having pulmonary diseases e.g. chronic bronchitis, cystic fibrosis or emphysema. The DNases are employed for enzymatic

alteration of the viscoelasticity of mucous, for treatment of patients with abnormal viscous, purulent secretions e.g. with infectious pneumonia, asthma, tuberculosis and fungal

infections. - In an example, purified deamidated human DNase and purified non-deamidated human DNase for use in this study were prepd. by

TCX chromatography. DNase enzymatic activity, synthetic double stranded DNA, 25 base pairs in length, was labelled with dinitrophenol (DNP) on one end and with biotin on the other end. Hydrolysis of the substrate by DNase was detected by capture of the reaction prods, on microtiter plate wells coated with antibody to DNP and by quantitation of the intact probe with streptavidin-horseradish peroxidase. Specific activity of stability samples was correlated (r power(2) = 0.613; n=5) with the extent of DNase

deamidation (range 27% - 93%). Extrapolation of the least squares linear equation provided an estimate that the specific activity of deamidated human DNase was approx. 77% lower than that of non-deamidated human DNase.

L5 ANSWER 57 OF 64 USPAT2 on STN ACCESSION NUMBER:

2005:189422 USPAT2 TITLE: Compaction assay for assessment of respiratory disease

therapy INVENTOR(S): Daugherty, Ann L., Palo Alto, CA, UNITED STATES Mrsny, Randy J., Redwood City, CA, UNITED STATES Patapoff, Thomas W., Belmont, CA, UNITED STATES

PATENT ASSIGNEE(S): Genentech, Inc., South San Francisco, CA, UNITED STATES (U.S. corporation)

	NUMBER	KIND	DATE	
PATENT INFORMATION: APPLICATION INFO.:	US 7193055 US 2005-33358	B2	20070320 20050110	(11)

RELATED APPLN. INPO.: Continuation of Ser. No. US 2002-162951, filed on 4 Jun 2002, ABABDONED Continuation of Ser. No. US 2001-771078, filed on 25 Jun 2001, ABABDONED Continuation of Ser. No. US 1997-840441, filed on 1 Apr 1997, ABABDONED Continuation of Ser. No. US 1997-840451, filed on 1 Apr 1997, ABABDONED Continuation of Ser. No. US

1995-539468, filed on 5 Oct 1995, ABANDONED Continuation of Ser. No. US 1994-355418, filed on 13 Dec 1994, ABANDONED Continuation of Ser. No. US 1993-132681, filed on 6 Oct 1993, ABANDONED

Continuation of Ser. No. US 1992-971019, filed on 2 Nov 1992, ARANDONED

DOCUMENT TYPE: Utility

FILE SEGMENT: GRANTED
PRIMARY EXAMINER: Carlson, Karen Cochrane

LEGAL REPRESENTATIVE: Evans, David W. NUMBER OF CLAIMS: 22

EXEMPLARY CLAIM: 1
NUMBER OF DRAWINGS: 9 Drawing Figure(s); 8 Drawing Page(s)
LINE COUNT: 1092

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A compaction assay measuring the viscoelasticity of sputum samples of patients subject to respiratory disease is provided. This assay is useful in determining the therapeutic efficacy of DNase, antibiotic and other respiratory disease treatments in improving lung function.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

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L5 ANSWER 54 OF 64 WPIDS COPYRIGHT 2009 THOMSON REUTERS on STN UADV USE/ADVANTAGE

Deamidated human DMase or non-deamidated human DMase may be used for treatment of patients having an accumulation of viscous, BNA-conty, material. Admin. of purified DMases pref. is via direct inhalation into the lungs. Non-deamidated human DMase may be admin. directly into the airway passages for the treatment of patients having pulmonary diseases e.g., chronic bronchitis, oyunic fibrosis or emphysems. The DMases are employed for enzymatic fibrosis or emphysems. The DMases are employed for enzymatic patients with abnormal viscous, purulent secretions e.g. with infectious pneumonia, asthma, tuberculosis and fungal infections.

In an example, purified deamidated human DNase and purified non-deamidated human DNase for use in this study were prepd. by.. . .

Member (0006)

ABEQ ZA 9303724 . . . and an immobilised non-hydrolysable DNA analogue

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Member (00:10)
AEGO DE 4392749 . . . and an immobilized non-hydrolysable DNA analogue resin.
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fibrosis or emphysems. The DMmars are employed for enzymatic alteration of the viscoelanticity of mucous, for treatment of patients with abnormal viscous, purulent secretions e.g. with infectious preumonia, asthma, tuberculosis and fungal infections. - In an example, purified desmidsted human DMmars or use in this study were prepd. ...

Member (GC11)

ABEQ JP 07507455 . . . and an immobilised non-hydrolysable DNA analogue

reain.

USE/ADVANTAGE - Deamidated human DNase or non-deamidated human DNase may be used for treatment of patients having an accumulation of direct inhalation into the lungs. Mon-deamidated human DNase may be admin. directly into the airway passages for the treatment of patients having pulsonary diseases e.g., chronic bronchlitis, cystic alteration of the viscoelasticity of smooth, pulson the patients of patients with shoreman viscoelasticity of smooth, pulson the patients of patients with shoreman viscoous, pursued sceretions e.g. with infectious pneumonia, author, tuberculosis and fungal under the patients with shoreman viscoous, pursued the ceretions e.g. with infectious pneumonia, author, tuberculosis and fungal under the patients with some distriction of the patients with some patients with some patients with some patients of the patients with some patients of the patients with some patients of the patients with the patients of the patients of the patients of the patients with the patients of the patie

Member (0018)

Member (0018) ABEQ US 5783433 . . and an immobilised non-hydrolysable DNA analogue resin.

DESCAUNATIAGE - Desmidated human UMase or non-deamidated human DMase may be used for treatment of patients having an accumulation of viscous, UMM-contg, materia; Admin of purified UMmases pref. is via direct inhalation into the image, Mon-deamidated human DMase may be admin. having pulsonary diseases e.g. chronic brocehitis, cystic fibrosis or emphysems. The UMmases are employed for enzymatic alteration of the viscoelasticity of mucous, for treatment of patients with abnormal viscous, pursuents exerctions e.g. with infectious pneumonis, astimas, tuberculosis and fungal image and purified non-deamidated human UMmase for use in this study were prept.

Member (0020)

ABEQ EP 1013284 . . . and an immobilised non-hydrolysable DNA analogue regin.

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alteration of the viscoelasticity of mucous, for treatment of
patients with abnormal viscous, purulent secretions e.g. with
infectious pneumonia, asthma, tuberculosis and fungal
infections. - In an example, purified desmi
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Member (0021)

ABBE EF 644932 . . and an immobilized non-hydrolysable DNA analogue resin.

USK-INDWATCME - Demaidated human DNaso or non-damidated human DNaso
may be used for treatment of patients having an accumulation of
viscous, DNA-cont, naterial. Admin. of purified DNasos part is via
direct inhalation into the lumps. Non-demaidated human DNaso may be admin.
directly into the airway passages for the treatment of patients
fibroais or employeem. The DNasos are employed for enzymatic
alteration of the viscoelasticity of musous, for treatment of
patients with abnormal viscous, purulent secretions e.g. with
infections permannia, atthms, tuberculosis and fungal
infections. - In an example, purified demaidated human DNaso and
purified non-demaidated human DNaso for use in this study were prepd. .

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